



Food and Drug Administration 10903 New Hampshire Avenue Document Control Center - WO66-G609 Silver Spring, MD 20993-0002

Mr. Robert F. Sullivan Senior Director FDA Regulatory Affairs STERIS Corporation 5960 Heisley Road Mentor, Ohio 44060

JAN 1 0 2017

Re: K092823

Trade/Device Name: Amsco Warming Cabinet

Regulation Number: 21 CFR 880.5725 Regulation Name: Infusion Pump

Regulatory Class: Class II

Product Code: LGZ

Dated: September 11, 2009 Received: September 14, 2009

Dear Mr. Sullivan:

This letter corrects our substantially equivalent letter of December 18, 2009.

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21)

CFR Part 807); labeling (21 CFR Parts 801 and 809); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

If you desire specific advice for your device on our labeling regulation (21 CFR Parts 801 and 809), please contact the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to

http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm.

Sincerely yours,

# Erin I. Keith -S

Erin I. Keith, M.S.
Director
Division of Anesthesiology,
General Hospital, Respiratory,
Infection Control, and Dental Devices
Office of Device Evaluation
Center for Devices and Radiological Health

Enclosure

## Indications for Use

510(k) Number (if known	ı):
Device Name:	Amsco Warming Cabinet
Indications For Use:	
sterile surgical irrigation :	binet is designed to raise the temperature of blankets, linens and solutions and IV solutions to an acceptable level for various gency, critical care and other healthcare applications
	•
Prescription Use(Part 21 CFR 801 Subpar	AND/OR Over-The-Counter Use X (21 CFR 801 Subpart C)
(PLEASE DO NOT WR PAGE IF NEEDED)	ITE BELOW THIS LINE-CONTINUE ON ANOTHER
	(Division Sign-Off) Division of Anesthesiology, General Hospital Infection Control, Dental Devices
	510(k) Number: <u>K-9 282 3</u>

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### 510(k) Summary For Amsco Warming Cabinet

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Summary Date:

September 11, 2009

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## STERIS TRADITIONAL 510(k) PREMARKET SUBMISSION STERIS Amsco Warming Cabinet

#### 1. Device Name

Trade Name:

**Amsco Warming Cabinet** 

Common/usual Name:

**Warming Cabinet** 

Classification Name:

Warmer, Irrigation Solution

Warmer, Cabinet

#### 2. Predicate Device

STERIS Amsco Warming Cabinet (Pre-Amendment)
Enthermics EC-7701 Fluid Warming Cabinet, K993797, January 20, 2000

#### 3. <u>Description of Device</u>

The Amsco Warming Cabinet is designed to store and warm sterile IV solutions, surgical irrigation solutions, linens and/or blankets to an acceptable level for hospital and surgical outpatient center applications.

The upper compartment of the 18" (457mm) deep model holds up to 24(1-liter) liquid bags or bottles and 14 (1-liter) IV solution bags; the lower compartment holds up to 72 (1-liter) liquid bags or bottles.

The upper compartment of the 24" (610mm) deep model holds up to 30 (1-liter) surgical flasks; the lower compartment holds up to 90 (1-liter) surgical flasks.

#### 4. Intended Use

The Amsco Warming Cabinet is designed to raise the temperature of blankets, linens and sterile surgical irrigation solutions and IV solutions to an acceptable level for various surgical, obstetrical, emergency, critical care and other healthcare applications.

### 5. <u>Description of Safety and Substantial Equivalence</u>

The Amsco Warming Cabinet is nearly identical to the two predicate devices in all material respects. A table comparing the technological characteristics of the proposed Amsco Warming Cabinet to the predicates is provided in Table 5-1.

Table 5-1 Summary of the Proposed Device and Predicate Devices
Technological Characteristics

	Amsco Warming Cabinet (IV Solution Capability)	Amsco Warming Cabinet	EC-7701 Fluid Warming
		(Pre-amendment)	Cabinet (K993797)
	The Amsco Warming Cabinet	Amsco Warming Cabinet is for	The Enthermics Medical
1 1 1	s designed to raise the	heating flasked solutions,	Systems EC-7701 Fluid
	emperature of blankets, linens	blankets and similar clinical	Warming Cabinet is
	and sterile surgical irrigation	articles.	designed to safely store
	solutions and IV solutions to	atticies.	and warm irrigation fluids
	an acceptable level for various		or injection fluids in
	surgical, obstetrical,		accordance with the
	emergency, critical care and		recommended warming
	other healthcare applications.		temperatures and storage
	The Amsco Warming Cabinet		times stated in the fluid
	s designed to hold a		manufacturer's labeling.
	combination of flasks and/or		manufacturer 3 taccting.
d	dry goods.		
	Electric heater and fan blower	Steam heat or Electric heater	Fully insulated
System (	(Convection heating)	blanket and fan blower	electrothermal cable array
		(Convection heating)	(Convection heating)
	Single/Double	Single/Dual	Single
Configuration			
	18" or 24"	18" or 24"	
	Wall or Counter	Wall or Counter	Wall
	Stainless Steel, ABS Plastic	Stainless Steel	Stainless Steel
Exterior a	and laminated galvanized steel		
Surfaces			
	ree-Standing (mobile) or	Open-Mounted or Recessed	Free-standing (mobile) or
1	Recessed		Recessed
Door S	Stainless Steel (Seliden)	God to the control of	
1 -	Stainless Steel (Solid and Glass)	Stainless Steel	Stainless Steel (Glass)
	8" upper / single chamber -	Dod Orace and 11	
	3.2 cu ft – up to 24 (1-liter)	Dual Compartment Model -	The cabinet is equipped
	oottles	Two shelves	with three (3) white,
Capacity	otties	15 flasks – 18" upper	epoxy-coated wire baskets,
1	8" lower chamber - 8.5 cu ft	45 flasks – 18" lower 20 flasks – 24" upper	each with a 24 liter
	- up to 72 (1-liter) bottles	60 flasks – 24" lower	capacity.
	up to 12 (1-incr) bottles	ou hasks = 24 lower	
2	4" upper / single chamber -	Single Compartment Model -	
4	1.3 cu ft – up to 30 (1-liter)	One shelf	
	pottles	15 flasks – 18"	1
		20 flasks – 24"	
2	24" lower chamber - 11.6 cu ft	_ ·	
J <i>-</i>	- up to 90 (1-liter) bottles		
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# STERIS TRADITIONAL 510(k) PREMARKET SUBMISSION STERIS Amsco Warming Cabinet

Features	Amsco Warming Cabinet	Amsco Warming Cabinet	EC-7701 Fluid Warming
· Garares	(IV Solution Capability)	(Pre-amendment)	Cabinet (K993797)
Cabinet	18" upper chamber - 3.1 cu ft	18" upper chamber - 3.1 cu ft	
Volume	24" upper chamber - 4.2 cu ft	24" upper chamber - 4.2 cu ft	•
	18" lower chamber - 8.9 cu ft	18" lower chamber - 8.9 cu ft	
<u> </u>	24" lower chamber - 12 cu ft	24" lower chamber - 12 cu ft	
Controls	Digital Push Button keypad /	Thermostat / power switch /	Electronic control consists
	power switch / Digital LCD	fuse with indicating light / color	of 4 digit LED display,
	temperature display / mode	coded temperature selector	on/off key, increase and
	selection buttons / door ajar		decrease keys, integrated
	indicator / Over-temperature		lock feature and a series of
	light for each compartment /		prompt sequence
	Data port for retrieval of stored temperatures.		indicators.
Software	Unit contains software	Not Applicable	Not applicable
Temperature	90°F (32°C) to 160°F (71°C)	95°F to 150°F (35°C to 65°C)	Not applicable 90°F (32°C) to 150°F
Selection	50 1 (52 C) to 100 1 (71 C)	(33 € 10 130 1 (33 € 10 03 €)	(66°C)
Range			(00 C)
Temperature	Temperature lock-out function	Not Available	The device allows the user
Lock	to prevent unauthorized		to "lock" the mode (IRR or
	temperature changes.		INJ) and temperature
			setting controls.
Door Lock	All configurations will be	Available by SSQ (Special	Available as an option
	equipped with either a manual	Sales Quote) only	-
	mechanical door lock or		
	optional electronic door lock		
Over	system for each compartment	 	
Temperature	Visual and audible alarm if unit has a chamber	Visual alarm if unit has a	Visual and audible alarm if
Alarm Point	_ · · · · · · · · · · · · · · · · · · ·	chamber temperature greater	unit has a chamber
Alain Foliit	temperature greater than 10°F (5.5°C) above set temperature.	than 12°F above set	temperature greater than
	In the event of an over temp	temperature. In the event of an	10°F (5.5°C) above set
	condition, sensors	over temp condition, sensors automatically turns off the	temperature. In the event
	automatically turns off the	heater(s).	of an over temp condition, the heating system shuts
	heater(s).	monter(3).	down.
Voltage	110/120 Vac, 220/240 Vac	Electric Model: 110/120 Vac,	125 Vac, 60 HZ, 1 ph
Requirements	nominal, 50/60 HZ	220/240 Vac nominal, 50/60	125 vac, 66 (12, 1 ph
-		HZ	
		Steam Model: 120 VAC single	
		phase	